**Kevin Doyle** 

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Evolutionary biologist with a strong passion for statistical data analysis and computational genomic research.

#### **Education**:

# **MA Biology - Applied Evolution**

Stony Brook University, Stony Brook, NY 2015

Relevant coursework: Biometry, Computational Biology, Population Genetics

GPA: 3.8

# **BS Biology - Ecology & Evolution**

Stony Brook University, Stony Brook, NY 2013

Minor: Ecosystems and Human Impact

# **Qualifications**:

 Unix Systems Algorithm Design

• R Git / GitHub

 Genomic Data Analysis Python

 HTML MS Office (Word, Excel)

Debugging / Troubleshooting

NGS Analysis

Software Compilation

Histology

#### Experience:

#### **Histology Technician**

Antech Diagnostics, Lake Success, NY

Nov 2016 - Present

Assisting in the histological process of identifying cancerous tumor progression for diagnostic purposes. Take thin sections around surgical margin of all diseased tissues and organs greater than 10 cm in length and fix them in paraffin.

- Use of a Laboratory Information System to input and manage data
- Take sections around surgical margin to determine extent of disease
- Work closely with fellow co-workers to meet daily quota minimums

## **Hollister Lab of Evolutionary Genomics**

Stony Brook University, Stony Brook, NY

Aug 2015 - May 2016

Project: Evolution of allopolyploidy in *Arabidopsis thaliana* 

Utilized Genome Analysis Toolkit (GATK) best practices pipeline to acquire, organize, and analyze Illumina next-generation sequencing (NGS) data from the National Center for Biotechnology Information (NCBI) Sequence Read Archive (SRA) database. Analyzed haplotype variance to infer global patterns of genomic divergence. Tools used were SRA Toolkit, SamTools, Picard Tools, Stampy, GATK, UnifiedGenotyper, HaplotypeCaller, VCFtools, SnpEff.

- Created bash scripts to enhance existing pipelines to streamline analyses
- Maintained data and documentation organized on Unix server
- · Monitored RAM and CPU usage from pipelines for maximum efficiency
- · Utilized GATK, Biopython, & R to analyze variation among individuals

# **Computational Biology Coursework**

Stony Brook University, Stony Brook, NY

Aug 2015 - Dec 2015

Created Python scripts to coincided with lectures pertaining to topics such as sequence alignment (pairwise & multiple), genome assembly, and gene finding using Hidden Markov Models. Course project involved creating functioning code from minimally existing Python code to perform an analysis on the accuracy of the duplication-mutation with complementarity (DMC) algorithm specified in *Navlakha & Kingsford 2011* on generating phylogenetic trees using a combination of protein-protein interaction (PPI) networks and genomic data.

# **Biometry Coursework**

Stony Brook University, Stony Brook, NY

Jan 2015 - May 2015

Rigorously utilized R to perform a wide variety of statistical analyses including linear/nonlinear regression, correlation, generalized linear models, probability distributions, maximum likelihood estimates, hypothesis testing, multiple comparisons analysis, bootstrapping and jackknifeing, data visualization with ggplot, ANOVA/ANCOVA, model selection/comparison, and principal component analysis.

# **Baines Lab of Aquatic Ecology**

Stony Brook University, Stony Brook, NY

Aug 2013 - May 2016

Project: Fe Limitation in the Copepod Acartia tonsa

Designed an experiment to determine the critical Fe:C threshold where iron limitation occurs in *Acartia tonsa* and performed statistical analyses using R and Excel to find where threshold occurred. Scrupulously clean conditions had to be maintained regarding all reagents and surfaces to prevent trace metal contamination.

- Maintained organized records of calculations and analyses
- Used R package 'segmented' to estimate where Fe threshold occurred by calculating the breakpoint between two separate linear regressions of egg production with replete Fe and egg production with deplete Fe
- Produced figures with R and Excel
- Grew cell and zooplankton cultures
- Created various chemical reagents to use in experiment
- Wrote paper (Estimating the Threshold of Iron Limitation in the Copepod Acartia tonsa) for publication

## Other Training:

## **Rosalind & Codecademy Exercises**

http://rosalind.info, http://codecademy.com

Using Python to complete problems under Rosalind's bioinformatics stronghold category and completing Codecademy course on SQL. Some topics completed thus far are:

- · Counting nucleotides and SNPs
- Transcribing and translating DNA
- Calculating probability of inherited alleles
- Dynamic programming
- · Finding DNA motifs